

# results of BLAST

#### BLASTP 2.2.6 [Apr-09-2003]

Reference:

Altschul, Stephen F., Thomas L. Madden, Alejandro A. Schäffer, Jinghui Zhang, Zheng Zhang, Webb Miller, and David J. Lipman (1997), "Gapped BLAST and PSI-BLAST: a new generation of protein database search programs", Nucleic Acids Res. 25:3389-3402.

RID: 1051730124-02802-29002

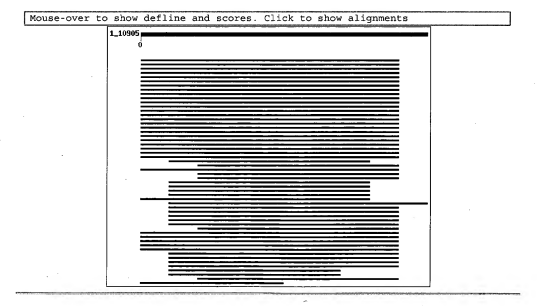
Query=

(10 letters)

If you have any problems or questions with the results of this search please refer to the  ${\tt BLAST\ FAQs}$ 

Taxonomy reports

### Distribution of 101 Blast Hits on the Query Sequence



Sequences producing significant alignments:

Score E (bits) Value

51730124-02802-29002,	http://www	http://www.ncbi.nlm.nih.gov/blast/Blast.cgi	
•			Bepalveir
			~ nouvair
gi 30048092 gb AAH50279.1  Similar to dynamin 1 [Homo sapiens]	35	0.12	ryerra
gi 539581 pir B40671 dynamin, internal form 2, short C-ter	35	0.12	2 4
gi 4758182 ref NP 004399.1 dynamin 1; dynamin; dynamin1; D	_35	0.12	
gi 181855 gb AAA02806.1  dynamin	35	0.12	9
gi 487853 gb AAA37319.1  dynamin	35	0.12	
gi 487855 gb AAA37323.1 dynamin	35	0.12	
gi   20824027   ref   XP   130133.1       dynamin [Mus musculus]       gi   539580   pir     A40671       dynamin, internal form 1, long C-term	_ <u>35</u> 35	0.12 0.12	
gi 18093102 ref NP 542420.1  dynamin 1 [Rattus norvegicus]	35	0.12	0
gi 21961254 gb AAH34679.1 Dnm protein [Mus musculus]	35	0.12	_
gi   181853   gb   AAA02805.1   dynamin	35	0.12	_
gi 729381 sp P39053 DYN1 MOUSE Dynamin-1 (Dynamin BREDNM19)	35	0.12	<u> </u>
gi 585074 sp Q08877 DYN3_RAT Dynamin 3 (Dynamin, testicular	31	2.3	9
gi 26331226 dbj BAC29343.1  unnamed protein product [Mus mu	31	2.3	
gi 24308091 ref NP 056384.1 KIAA0820 protein [Homo sapiens gi 28972415 dbj BAC65661.1 mKIAA0820 protein [Mus musculus]	31 31	2.3	
gi 27805466 sp Q9UQ16 DYN3 HUMAN Dynamin 3 (Dynamin, testic	31	2.3	
gi 1835967 gb AAB46800.1 dynamin III isoform=GTPase homolo	31	2.3	
gi   19924077   ref   NP 612547.1   testicular dynamin [Rattus nor	31	2.3	
gi 19353648 gb AAH24584.1 9630020E24Rik protein [Mus muscu	31	2.3	_
gi 27369922 ref NP 766234.1 RIKEN cDNA 9630020E24 [Mus mus	<u>31</u>	2.3	
gi 20521666 dbj BAA74843.2 KIAA0820 protein [Homo sapiens]	31	2.3	E
gi 21288051 gb EAA00372.1  agCP9929 [Anopheles gambiae str	25	139	m
gi   17540448   ref   NP_501236.1   Predicted CDS, putative nuclea gi   22988735   ref   ZP_00033796.1   hypothetical protein [Burkho	<u>25</u> 25	139 139	
gi 27378665 ref NP_770194.1 blr3554 [Bradyrhizobium japoni	25	139	4
gi 24474936 emb CAC87573.1 trypanothione synthetase [Trypa	24	186	
gi   17570421   ref   NP 509982.1   Putative plasma membrane membr	24	186	
gi 29468074 gb AA000721.1 trypanothione synthetase [Trypan	24	186	
gi 24655701 ref NP 523793.2 Focal adhesion kinase-like CG1	24	250	
gi   6525023   gb   AAF15292.1   AF201701 1 focal adhesion kinase h	24	250	L L
gi   6409130   gb   AAF07854.1   AF112116   focal adhesion kinase h gi   6016830   dbj   BAA85188.1   focal adhesion kinase [Drosophil	$\frac{24}{24}$	250 250	
gi 21289343 gb EAA01636.1 ebiP7766 [Anopheles gambiae str	$\frac{24}{24}$	335	
gi 7657238 ref NP 055240.1  inversin [Homo sapiens] >gi 392	24	335	0
gi 28372796 gb AAO18093.1 unknown [Babesia microti]	24	335	
gi 15237241 ref NP_200085.1  hypothetical protein; protein	24	335	
gi 3925425 gb AAC79457.1 inversin protein alternative isof	24	335	
gi   14574596   gb   AAD02131.2   inv candidate homolog [Homo sapi gi   6912374   ref   NP 036317.1   forkhead box E2 [Homo sapiens]	$\frac{24}{24}$	335	2
gi   6912374   ref   NP   036317.1         forkhead box E2 [Homo sapiens]         gi   21222187   ref   NP   627966.1         putative membrane protein [Str	$\frac{24}{24}$	335 335	
gi 6978771 ref NP 037331.1  dynamin 2 [Rattus norvegicus] >	23	450	<b>.</b>
gi 27806155 ref NP_776893.1  synaptojanin 1 [Bos taurus] >g	23	450	0
gi   12836637   dbj   BAB23745.1   unnamed protein product [Mus mu	23	450	
gi 2702321 gb AAC51921.1  synaptojanin [Homo sapiens]	23	450	
gi 15609426 ref NP 216805.1 cdh [Mycobacterium tuberculosi	23	450	
gi 28919283   gb EAA28749.1   predicted protein [Neurospora cr	23	450	
gi 29732208 ref   XP_295554.1   hypothetical protein XP_295554   gi 10720298 sp   018964   SYJ1_BOVIN Synaptojanin 1 (Synaptic i	23	450 450	
gi   16307381   gb   AAH10233.1   Dnm2 protein [Mus musculus]	23	450	10
gi 9625476 ref NP_039710.1 predicted 16.3kd protein [Mycob	23	450	_
gi   6681207   ref   NP 031897.1   dynamin 2 [Mus musculus] >gi   48	23	450	
gi 20521692 dbj BAA74933.2  KIAA0910 protein [Homo sapiens]	23	450	
<u>gi 4507335 ref NP 003886.1 </u> synaptojanin 1; inositol 5'-pho	23	450	
gi 22002044 sp P39054 DYN2 MOUSE Dynamin 2 (Dynamin UDNM)	_23	450	
gi 6225895 sp P56722 PTPN BOVIN Protein-tyrosine phosphatas	_23	450	
gi   24586224   ref   NP   724552.1   CG11112-PB   [Drosophila melanog		450	
gi 1363285 pir  A36878 dynamin 2 - rat >gi 404073 gb AAA167	_23	450	

gi 1083647 pir  B53165 dynamin II isoform ba - rat	23	450	
gi 24644378 ref NP 730989.1 pollux CG1093-PB >gi 23170404	23	604	
gi 29727084 ref XP 298121.1  hypothetical protein XP 298121	23	604	L
qi 1362003 pir   S55889 protein phosphatase 2A B regulatory	23	604	
gi 25453269 sp Q9GLG4 SG1 PIG Secretogranin I precursor (Sg	23	604	
gi   11034601   dbj   BAB17125.1   P0463F06.12 [Oryza sativa (japo		604	
gi 19112873 ref NP 596081.1 probable serine threonine-prot	23	604	
gi 22507080 gb AAM97755.1 hypothetical protein [Oryza sati	23	604	L
gi 21427013   gb   AAM53039.1   AF262024 1 PJA1 [Homo sapiens]	<u>23</u>	604	15
gi 23103921 ref ZP 00090393.1 hypothetical protein [Azotob gi 11290035 pir T48835 lethal(2)denticleless related prote	- <u>23</u> - 23	604 604	
gi 23103819 ref ZP 00090293.1 hypothetical protein [Azotob	23	604	
gi   13195576   gb   AAK15765.1   AF335251 1 Prajal isoform c [Mus	23	604	
gi 23471390 ref ZP 00126720.1 hypothetical protein [Pseudo	23	604	_
gi 29841092 gb AAP06105.1 similar to GenBank Accession Num	23	604	
gi 22968437 gb ZP_00016025.1 hypothetical protein [Rhodosp	23	604	
gi 26381879 dbj BAC25475.1 unnamed protein product [Mus mu	23	604	L
gi 18403637 ref NP_564595.1 55 kDa B regulatory subunit of	23	604	
gi 21356709 ref NP_650112.1  CG6923-PA [Drosophila melanoga	23	604	L
gi   12837873   dbj   BAB23982.1   unnamed protein product [Mus mu	23	604	L
gi 13195574 qb AAK15764.1 AF335250 1 Prajal isoform a [Mus	23	604	
qi 21295549 qb EAA07694.1 agCP1713 [Anopheles gambiae str	23	604	
gi 11283372 pir T48801 hypothetical protein 15E6.90 [impor	23	604	
gi 9629417 ref NP 044638.1 very large tegument protein [Hu	_23	604	
gi 19113573 ref NP 596781.1 similarity to yeast suppressor	23	604	
gi 25405491 pir   G96555 55 kDa B regulatory subunit of phos	23	604	
gi         22972926         gb         ZP         00019777.1         hypothetical protein [Chlorof           qi         22902385         gb         AAH37616.1         Pjal protein [Mus musculus]	23	604 604	
<del>#                                </del>		604	L.
gi 7511991 pir  T13718 pollux gene protein - fruit fly (Dro gi 22507073 gb AAM97748.1  hypothetical protein [Oryza sati	<u>23</u>	604	-
<del>2     2  </del>	23	604	
gi   1373163   gb   AAB02200.1   pollux gi   23063477   ref   ZP 00088208.1   hypothetical protein [Pseudo	23	604	
gi 23060056 ref ZP_00084981.1 hypothetical protein [Pseudo	23	604	
gi 20888961 ref XP 146899.1  similar to circumsporozoite pr	23	604	L
qi 19112067 ref NP 595275.1 26s protease subunit [Schizosa	23	604	_
gi 7492202 pir T42085 MPT4 protein homolog - fission yeast	23	604	
gi 5305335 gb AAD41594.1 AF071081 1 proline-rich mucin homo	23	604	
gi 20838555 ref XP_162177.1  hypothetical protein XP_162177	23	604	L.
gi 24644376 ref NP 730988.1  pollux CG1093-PA >gi 7296688 g	23	604	
gi 12831443 gb AAK08598.1 unknown [Agrobacterium tumefaciens]	22	810	
gi 29738779 ref XP_295680.1  hypothetical protein XP_295680	22	810	
gi 22093838 dbj BAC07125.1 hypothetical protein~predicted	22	810′	

#### Alignments

Deselect all

Select all

Get selected sequences

```
□ >gi | 30048092 | gb | AAH50279.1 | Similar to dynamin 1 [Homo sapiens]

Length = 881

Score = 35.0 bits (75), Expect = 0.12
Identities = 10/10 (100%), Positives = 10/10 (100%)

Query: 1 QVPSRPNRAP 10
QVPSRPNRAP
Sbjct: 861 QVPSRPNRAP 870

□ >gi | 539581 | pir | | B40671 | dynamin, internal form 2, short C-terminal form - human
Length = 851
```

```
Score = 35.0 bits (75), Expect = 0.12
 Identities = 10/10 (100%), Positives = 10/10 (100%)
          QVPSRPNRAP 10
Query: 1
           OVPSRPNRAP
Sbjct: 831 QVPSRPNRAP 840
                                 dynamin 1; dynamin; dynamin1; Dynamin-1 [Homo sapi
3 >qi | 4758182 | ref | NP 004399.1
 gi | 461976 | sp | Q05193 | DYN1 HUMAN
                                  Dynamin-1
 gi | 181849 | gb | AAA02803.1 |
                             dynamin
         Length = 864
 Score = 35.0 \text{ bits } (75), Expect = 0.12
 Identities = 10/10 (100%), Positives = 10/10 (100%)
Query: 1 QVPSRPNRAP 10
           QVPSRPNRAP
Sbjct: 831 QVPSRPNRAP 840
dynamin
         Length = 126
 Score = 35.0 bits (75), Expect = 0.12
 Identities = 10/10 (100%), Positives = 10/10 (100%)
          QVPSRPNRAP 10
Query: 1
           OVPSRPNRAP
Sbjct: 106 QVPSRPNRAP 115
dynamin
          Length = 243
 Score = 35.0 bits (75), Expect = 0.12
 Identities = 10/10 (100%), Positives = 10/10 (100%)
Query: 1
          QVPSRPNRAP 10
           QVPSRPNRAP
Sbjct: 210 QVPSRPNRAP 219
____>gi|487855|gb|AAA37323.1|
                               dynamin
         Length = 612
 Score = 35.0 bits (75), Expect = 0.12
 Identities = 10/10 (100%), Positives = 10/10 (100%)
Query: 1
          QVPSRPNRAP 10
          QVPSRPNRAP
Sbjct: 592 QVPSRPNRAP 601
| >gi | 20824027 | ref | XP | 130133.1 | dynamin [Mus musculus]
         Length = 864
 Score = 35.0 bits (75), Expect = 0.12
 Identities = 10/10 (100%), Positives = 10/10 (100%)
          OVPSRPNRAP 10
Query: 1
          OVPSRPNRAP
Sbjct: 827 QVPSRPNRAP 836
□ >gi|539580|pir||A40671
                           dynamin, internal form 1, long C-terminal form - human
         Length = 864
Score = 35.0 bits (75), Expect = 0.12
```

```
Identities = 10/10 (100\%), Positives = 10/10 (100\%)
         QVPSRPNRAP 10
Query: 1
            QVPSRPNRAP
Sbjct: 831 QVPSRPNRAP 840
| >gi | 18093102 | ref | NP 542420.1 | dynamin 1 [Rattus norvegicus] | gi | 118966 | sp | P21575 | DYN1 RAT | Dynamin-1 (D100) (Dynamin, brain) (B-dynamin)
 gi 111573 pir | S11508
                         D100 protein - rat
                              D100 [Rattus norvegicus]
 gi | 56054 | emb | CAA38397.1 |
 gi 227123 prf | 1614348A
                            dynamin 1 D100 protein
          Length = 851
 Score = 35.0 bits (75), Expect = 0.12
 Identities = 10/10 (100%), Positives = 10/10 (100%)
           QVPSRPNRAP 10
Ouery: 1
           OVPSRPNRAP
Sbjct: 831 QVPSRPNRAP 840
___>gi|21961254|gb|AAH34679.1|
                                  Dnm protein [Mus musculus]
          Length = 867
 Score = 35.0 bits (75), Expect = 0.12
 Identities = 10/10 (100%), Positives = 10/10 (100%)
Query: 1
           QVPSRPNRAP 10
            QVPSRPNRAP
Sbjct: 831 QVPSRPNRAP 840
dynamin
          Length = 131
 Score = 35.0 \text{ bits } (75), \text{ Expect = } 0.12
 Identities = 10/10 (100%), Positives = 10/10 (100%)
Query: 1
           QVPSRPNRAP 10
           OVPSRPNRAP
Sbjct: 106 QVPSRPNRAP 115
___ >gi | 729381 | sp | P39053 | DYN1 MOUSE
                                        Dynamin-1 (Dynamin BREDNM19)
qi|487857|qb|AAA37324.1|
                              dynamin
          Length = 861
 Score = 35.0 bits (75), Expect = 0.12
 Identities = 10/10 (100%), Positives = 10/10 (100%)
Query: 1
           QVPSRPNRAP 10
           QVPSRPNRAP
Sbjct: 827 QVPSRPNRAP 836
□>gi|585074|sp|Q08877|DYN3 RAT Dynamin 3 (Dynamin, testicular) (T-dynamin)
gi 2143992 pir | I55498 testicular dynamin - rat
 gi | 391872 | dbj | BAA03161.1 |
                               testicular dynamin [Rattus norvegicus]
         Length = 848
 Score = 30.8 bits (65), Expect = 2.3
 Identities = 9/10 (90%), Positives = 9/10 (90%)
           OVPSRPNRAP 10
Query: 1
           QVPSRP RAP
Sbjct: 823 QVPSRPTRAP 832
```

```
unnamed protein product [Mus musculus]
| >qi | 26331226 | dbj | BAC29343.1 |
          Length = 863
 Score = 30.8 bits (65), Expect = 2.3
 Identities = 9/10 (90%), Positives = 9/10 (90%)
          QVPSRPNRAP 10
Query: 1
           QVPSRP RAP
Sbjct: 827 QVPSRPTRAP 836
__|>gi|24308091|ref|NP 056384.1|
                                 KIAA0820 protein [Homo sapiens]
 gi | 12052944 | emb | CAB66647.1 |
                              hypothetical protein [Homo sapiens]
          Length = 863
 Score = 30.8 bits (65), Expect = 2.3
 Identities = 9/10 (90%), Positives = 9/10 (90%)
Query: 1 QVPSRPNRAP 10
          OVPSRP RAP
Sbjct: 827 QVPSRPTRAP 836
Sqi | 28972415 | dbj | BAC65661.1 | mKIAA0820 protein [Mus musculus]
         Length = 452
 Score = 30.8 bits (65), Expect = 2.3
 Identities = 9/10 (90%), Positives = 9/10 (90%)
Query: 1
          QVPSRPNRAP 10
          OVPSRP RAP
Sbjct: 416 QVPSRPTRAP 425
___>gi|27805466|sp|Q9UQ16|DYN3_HUMAN
                                      Dynamin 3 (Dynamin, testicular) (T-dynamin)
         Length = 859
 Score = 30.8 bits (65), Expect = 2.3
 Identities = 9/10 (90%), Positives = 9/10 (90%)
Query: 1 QVPSRPNRAP 10
          OVPSRP RAP
Sbjct: 823 QVPSRPTRAP 832
spliced} [rats, Sprague-Dawley, brain, Peptide Partial,
         42 aa]
         Length = 42
 Score = 30.8 bits (65), Expect = 2.3
 Identities = 9/10 (90%), Positives = 9/10 (90%)
Query: 1 QVPSRPNRAP 10
         QVPSRP RAP
Sbjct: 6 QVPSRPTRAP 15
□>gi|19924077|ref|NP_612547.1| testicular dynamin [Rattus norvegicus]
gi|6409115|gb|AAF07848.1|AF201839 1
                                    dynamin IIIbb isoform [Rattus norvegicus]
         Length = 869
 Score = 30.8 bits (65), Expect = 2.3
 Identities = 9/10 (90%), Positives = 9/10 (90%)
          OVPSRPNRAP 10
Query: 1
          QVPSRP RAP
Sbjct: 833 QVPSRPTRAP 842
```

```
__>gi|19353648|gb|AAH24584.1| 9630020E24Rik protein [Mus musculus]
          Length = 319
 Score = 30.8 \text{ bits } (65), \text{ Expect = } 2.3
 Identities = 9/10 (90\%), Positives = 9/10 (90\%)
          QVPSRPNRAP 10
Query: 1
           OVPSRP RAP
Sbjct: 294 QVPSRPTRAP 303
☐>gi|27369922|ref|NP 766234.1|
                                  RIKEN cDNA 9630020E24 [Mus musculus]
 gi | 26340464 | dbj | BAC33895.1 |
                               unnamed protein product [Mus musculus]
          Length = 859
 Score = 30.8 bits (65), Expect = 2.3
 Identities = 9/10 (90%), Positives = 9/10 (90%)
Query: 1
          OVPSRPNRAP 10
           QVPSRP RAP
Sbjct: 823 QVPSRPTRAP 832
____>qi|20521666|dbj|BAA74843.2|
                                 KIAA0820 protein [Homo sapiens]
        Length = 892
 Score = 30.8 bits (65), Expect = 2.3
 Identities = 9/10 (90%), Positives = 9/10 (90%)
          OVPSRPNRAP 10
Query: 1
           QVPSRP RAP
Sbjct: 856 QVPSRPTRAP 865
>qi|21288051|qb|EAA00372.1|. aqCP9929 [Anopheles gambiae str. PEST]
          Length = 332
 Score = 24.8 bits (51), Expect = 139
 Identities = 8/13 (61%), Positives = 10/13 (76%), Gaps = 3/13 (23%)
Query: 1 QVPSRP---NRAP 10
          QVP+RP
                   +RAP
Sbjct: 180 QVPNRPAPVDRAP 192
>gi|17540448|ref|NP 501236.1
                                   Predicted CDS, putative nuclear protein family me
            specific [Caenorhabditis elegans]
 gi|7503465|pir||T25752 hypothetical protein F45E4.4 - Caenorhabditis elegans
 gi 13559603 gb AAK29815.1 Hypothetical protein F45E4.4 [Caenorhabditis elegans]
         Length = 2361
 Score = 24.8 bits (51), Expect = 139
 Identities = 8/11 (72%), Positives = 10/11 (90%), Gaps = 1/11 (9%)
           QVPSR-PNRAP 10
Query: 1
           QVPSR P+R+P
Sbjct: 1405 QVPSRQPSRSP 1415
□>gi|22988735|ref|ZP 00033796.1| hypothetical protein [Burkholderia fungorum]
         Length = 412
 Score = 24.8 bits (51), Expect =
                                  139
 Identities = 7/8 (87%), Positives = 8/8 (100%)
Query: 2
          VPSRPNRA 9
          VPSRP+RA
```

```
Sbjct: 111 VPSRPSRA 118
p; | 27378665 | ref | NP 770194.1 | blr3554 [Bradyrhizobium japonicum]
 gi 27351814 dbj BAC48819.1
                               blr3554 [Bradyrhizobium japonicum USDA 110]
          Length = 136
 Score = 24.8 bits (51), Expect = 139
 Identities = 7/8 (87%), Positives = 7/8 (87%)
Query: 3 PSRPNRAP 10
          PSRPNR P
Sbjct: 43 PSRPNRDP 50
□| >qi | 24474936 | emb | CAC87573.1 | trypanothione synthetase [Trypanosoma brucei]
          Length = 627
 Score = 24.4 bits (50), Expect =
                                    186
 Identities = 7/8 (87%), Positives = 7/8 (87%)
           PSRPNRAP 10
Query: 3
           PSRPNR P
Sbjct: 189 PSRPNRNP 196
| >gi | 17570421 | ref | NP_509982.1 |
                                    Putative plasma membrane membrane protein, with a
           transmembrane domains, nematode specific [Caenorhabditis
           elegans]
 gi | 7510333 | pir | | T27257
                         hypothetical protein Y62H9A.1 - Caenorhabditis elegans
 gi 3881218 emb CAA21558.1 Hypothetical protein Y62H9A.1 [Caenorhabditis elegans]
          Length = 366
 Score = 24.4 bits (50), Expect =
                                     186
 Identities = 8/11 (72%), Positives = 8/11 (72%), Gaps = 1/11 (9%)
Query: 1
           QVPSRP-NRAP 10
            VPSRP NR P
Sbjct: 349 HVPSRPTNRVP 359
>gi|29468074|gb|AA000721.1|
                                 trypanothione synthetase [Trypanosoma brucei brucei]
          Length = 627
 Score = 24.4 bits (50), Expect =
 Identities = 7/8 (87%), Positives = 7/8 (87%)
Query: 3
           PSRPNRAP 10
           PSRPNR P
Sbjct: 189 PSRPNRNP 196
sgi|24655701|ref|NP_523793.2|
                                     Focal adhesion kinase-like CG10023-PA
gi 24655706 ref NP 725891.1
                                  Focal adhesion kinase-like CG10023-PB
 gi 24655711 ref NP_725892.1
                                 Focal adhesion kinase-like CG10023-PC
gi | 7302478 | gb | AAF57562.1 |
                               CG10023-PA [Drosophila melanogaster]
 gi 21645172 gb AAM70852.1
                                CG10023-PB [Drosophila melanogaster]
 gi | 21645173 | gb | AAM70853.1 |
                                CG10023-PC [Drosophila melanogaster]
          Length = 1200
 Score = 24.0 bits (49), Expect =
                                    250
 Identities = 8/9 (88%), Positives = 8/9 (88%), Gaps = 1/9 (11%)
Query: 2
           VPSR-PNRA 9
           VPSR PNRA
Sbjct: 943 VPSRPPNRA 951
| >gi | 6525023 | gb | AAF15292.1 | AF201701 1
                                            focal adhesion kinase homolog FAK56 [Droso
```

```
Length = 1200
 Score = 24.0 bits (49), Expect =
                                  250
 Identities = 8/9 (88%), Positives = 8/9 (88%), Gaps = 1/9 (11%)
           VPSR-PNRA 9
Query: 2
           VPSR PNRA
Sbjct: 943 VPSRPPNRA 951
__>gi|6409130|gb|AAF07854.1|AF112116_1 focal adhesion kinase homolog DFak56 [Dros
          Length = 1200
 Score = 24.0 bits (49), Expect =
                                   250
 Identities = 8/9 (88%), Positives = 8/9 (88%), Gaps = 1/9 (11%)
Query: 2
           VPSR-PNRA 9
           VPSR PNRA
Sbjct: 943 VPSRPPNRA 951
- sgi|6016830|dbj|BAA85188.1| focal adhesion kinase [Drosophila melanogaster]
          Length = 1198
 Score = 24.0 bits (49), Expect =
                                   250
 Identities = 8/9 (88%), Positives = 8/9 (88%), Gaps = 1/9 (11%)
Query: 2 VPSR-PNRA 9
           VPSR PNRA
Sbjct: 941 VPSRPPNRA 949
\square >gi|21289343|gb|EAA01636.1| ebiP7766 [Anopheles gambiae str. PEST]
          Length = 459
 Score = 23.5 bits (48), Expect =
                                  335
 Identities = 9/19 (47%), Positives = 9/19 (47%), Gaps = 10/19 (52%)
Query: 2
          VPSRPN-----RAP 10
           VPSRPN
                           PAP
Sbjct: 208 VPSRPNSSGGRTTDASRAP 226
___>gi|765723<u>8|ref|NP_055240.1|</u>
                                inversin [Homo sapiens]
 gi|3925387|gb|AAC79436.1| inversin protein [Homo sapiens]
 gi|3925424|qb|AAC79456.1|
                             inversin protein [Homo sapiens]
          Length = 1065
 Score = 23.5 bits (48), Expect = 335
 Identities = 7/9 (77%), Positives = 8/9 (88%)
Query: 2
          VPSRPNRAP 10
           VPSR +RAP
Sbjct: 632 VPSRQSRAP 640
sgi|28372796|gb|AAO18093.1| unknown [Babesia microti]
         Length = 253
 Score = 23.5 bits (48), Expect =
                                  335
 Identities = 7/9 (77%), Positives = 8/9 (88%)
Query: 2 VPSRPNRAP 10
          VPSRP+ AP
Sbjct: 88 VPSRPHSAP 96
Sgi|15237241|ref|NP_200085.1| hypothetical protein; protein id: At5g52730.1 [Ara
           thaliana]
```

```
gi | 8953728 | dbj | BAA98091.1 |
                              emb|CAA71173.1~gene id:F6N7.22~similar to unknown prot
           [Arabidopsis thaliana]
          Length = 185
 Score = 23.5 bits (48), Expect = 335
 Identities = 8/11 (72%), Positives = 8/11 (72%), Gaps = 3/11 (27%)
           PSR---PNRAP 10
Query: 3
           PSR
                 PNRAP
Sbjct: 140 PSREPEPNRAP 150
∐>gi|3925425|gb|AAC79457.1| inversin protein alternative isoform [Homo sapiens]
          Length = 895
 Score = 23.5 bits (48), Expect = 335
 Identities = 7/9 (77%), Positives = 8/9 (88%)
Query: 2
           VPSRPNRAP 10
           VPSR +RAP
Sbjct: 632 VPSRQSRAP 640
sqi|14574596|gb|AAD02131.2| inv candidate homolog [Homo sapiens]
          Length = 1013
 Score = 23.5 bits (48), Expect =
                                    335
 Identities = 7/9 (77%), Positives = 8/9 (88%)
           VPSRPNRAP 10
Query: 2
           VPSR +RAP
Sbjct: 580 VPSRQSRAP 588
____>gi|6912374|ref|NP 036317.1|
                                   forkhead box E2 [Homo sapiens]
 gi | 20177899 | sp | Q99526 | FXE2_HUMAN
                                      Forkhead box protein E2 (HNF-3/fork head-like p
           (HFKH4)
 gi | 1770432 | emb | CAA64246.1 |
                                fork head like protein [Homo sapiens]
          Length = 500
 Score = 23.5 bits (48), Expect =
                                    335
 Identities = 8/11 (72%), Positives = 8/11 (72%), Gaps = 2/11 (18%)
Query: 2
           VPSR--PNRAP 10
           VP R PNRAP
Sbjct: 323 VPPRAPPNRAP 333
☐ >gi|21222187|ref|NP_627966.1| putative membrane protein [Streptomyces coelicolor
 gi | 10432480 | emb | CAC10314.1 | putative membrane protein [Streptomyces coelicolor A3
          Length = 180
 Score = 23.5 bits (48), Expect = 335
 Identities = 7/9 (77%), Positives = 8/9 (88%)
Query: 1 QVPSRPNRA 9
          Q P+RPNRA
Sbjct: 6 QPPARPNRA 14
□>gi|6978771|ref|NP_037331.1|
                                  dynamin 2 [Rattus norvegicus]
 gi 729380 sp P39052 DYN2 RAT
                                Dynamin 2
 gi|1083646|pir||A53165 dynamin II isoform aa - rat
gi | 416396 | gb | AAA19736.1 |
                              dynamin IIaa
          Length = 870
 Score = 23.1 bits (47), Expect =
 Identities = 7/10 (70%), Positives = 8/10 (80%)
```

## chimera (ki-mer'a, ki-)

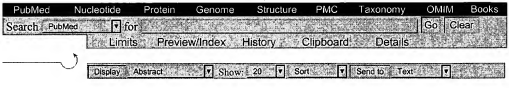
- In experimental embryology, the individual produced by grafting an embryonic part of one animal on to the embryo of another, either of the same or of another species.
- 2. An organism that has received a transplant of genetically and immunologically different tissue, such as bone marrow.
- 3. Dizygotic twins that retain each other as immunologically distinct types of erythrocytes.
- 4. A protein fusion in which two different proteins are linked via peptide bonds; usually genetically engineered. Chimeric antibodies may have the Fab fragment from one species fused with the Fc fragment from another.
- Any macromolecule fusion formed by two or more macromolecules from different species or from different genes.
- [L. Chimaera, G. Chimaira, mythic monster (lit. a she-goat)]











☐1: Biol Chem 2003 Feb;384(2):193-202

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Thrombin signaling in the brain: the role of protease-activated receptors.

Wang H, Reiser G.

PubMed Services Institut fur Neurobiochemie, Medizinische Fakultat der Otto-von-Guericke-Universitat Magdeburg, Leipziger Strasse 44, D-39120 Magdeburg, Germany.

Signaling by the protease thrombin has started to be appreciated in cell biology, especially since the gene for protease-activated receptor-1 (PAR-1) has been cloned. Apart from the central role of thrombin in blood coagulation and wound healing, thrombin also regulates cellular functions in a large variety of cells through PAR-1, PAR-3 and PAR-4. Receptors are activated by a proteolytic cleavage mechanism via G protein-coupled signaling pathways. Accumulating evidence shows that thrombin changes the morphology of neurons and astrocytes, induces glial cell proliferation, and even exerts, depending on the concentration applied, either cytoprotective or cytotoxic effects on neural cells. These effects may be mediated, through either distinct or overlapping signal transduction cascades, by activation of PARs. This review focuses on the underlying signaling events initiated by thrombin in neuronal and glial cells, to summarize our understanding of the intracellular signaling machinery linking thrombin receptors to their potential physiological and pathological functions in the CNS.

Related Resources

PMID: 12675511 [PubMed - in process]

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